

# The molecular phylogenetic position of Harpagocarpus (Polygonaceae) sheds new light on the infrageneric classification of Fagopyrum

Daozhang Min<sup>1,2,3\*</sup>, Wei Shi<sup>1\*</sup>, Mohammad Mehdi Dehshiri<sup>4\*</sup>, Yuting Gou<sup>2</sup>, Wei Li<sup>5</sup>, Kaixuan Zhang<sup>5</sup>, Meiliang Zhou<sup>5</sup>, Bo Li<sup>2,3</sup>

I State Key Laboratory of Desert and Oasis Ecology, Xinjiang Institute of Ecology and Geography, Chinese Academy of Sciences, Urumqi 830011, China 2 Research Centre of Ecological Sciences, College of Agronomy, Jiangxi Agricultural University, Nanchang 330045, China 3 The Specimen Museum of Xinjiang Institute of Ecology and Geography, Chinese Academy of Sciences, Urumqi 830011, China 4 Department of Biology, Borujerd Branch, Islamic Azad University, Borujerd, Iran 5 Institute of Crop Sciences, Chinese Academy of Agricultural Sciences, Beijing 100081, China

Corresponding authors: Meiliang Zhou (zhoumeiliang@caas.cn), Bo Li (hanbolijx@163.com)

Academic editor: A. Sukhorukov | Received 17 November 2022 | Accepted 19 January 2023 | Published 28 February 2023

**Citation:** Min D, Shi W, Dehshiri MM, Gou Y, Li W, Zhang K, Zhou M, Li B (2023) The molecular phylogenetic position of *Harpagocarpus* (Polygonaceae) sheds new light on the infrageneric classification of *Fagopyrum*. PhytoKeys 220: 109–126. https://doi.org/10.3897/phytokeys.220.97667

#### **Abstract**

In the context of the molecular phylogeny of Polygonaceae, the phylogenetic positions of most genera and their relationships have been resolved. However, the monotypic genus *Harpagocarpus* has never been included in any published molecular phylogenetic studies. In the present study, we adopt a two-step approach to confirm the phylogenetic placement of *Harpagocarpus* using two datasets: (1) a concatenated dataset of three chloroplast DNA (cpDNA) regions (*matK*, *rbcL* and *trnL-F*) for Polygonaceae and (2) a combined cpDNA dataset of five sequences (*accD*, *matK*, *psbA-trnH*, *rbcL* and *trnL-F*) for *Fagopyrum*. Our analyses confirm the previous hypothesis based on morphological, anatomical and palynological investigations that *Harpagocarpus* is congeneric with *Fagopyrum* and further reveal that *H. snowdenii* ( $\equiv E$  *snowdenii*) is sister to the woody buckwheat *E tibeticum*. Within *Fagopyrum*, three highly supported clades were discovered and the first sectional classification was proposed to accommodate them: sect. *Fagopyrum* comprises the two domesticated common buckwheat (*F. esculentum* and *F. tataricum*) and their wild relatives (*F. esculentum* subsp. *ancestrale*, *F. homotropicum* and *F. dibotrys*) which are characterised by having large corymbose inflorescences and achenes greatly exceeding the perianth; sect. *Tibeticum*, including *E. snowdenii* and *E. tibeticum*, is characterised by the achene having appurtenances along the ribs, greatly exceeding the perianth and the

These authors contributed equally to this work.

perianth accrescent in fruit; sect. *Urophyllum* contains all other species of which the achenes were completely enclosed in the perianth. This study is very helpful to understand the phylogeny of the *Fagopyrum* and sheds light on the future study of taxonomy, biogeography, diversification and character evolution of the genus.

#### **Keywords**

buckwheat, cpDNA, Fagopyreae, morphology, new section

#### Introduction

Polygonaceae, a family of the flowering plants known as the buckwheat family, can be easily distinguished by its ocrea, orthotropous ovules, trigonal (typically) achenes and quincuncial aestivation (Judd et al. 2007) and is found in almost all ecosystems (Sanchez et al. 2009). Numerous molecular phylogenetic analyses (e.g. Cuénoud et al. (2002); Schäferhoff et al. (2009); Moore et al. (2010); Yang et al. (2015); Walker et al. (2018); Yao et al. (2019); Li et al. (2021)) have provided strong evidence for the monophyly of Polygonaceae and the family's membership in the FTPP clade of the order Caryophyllales, which also includes the Plumbaginaceae, Polygonaceae, Tamaricaceae and Frankeniaceae, has been securely supported (e.g. Cuénoud et al. (2002); Brockington et al. (2009); Walker et al. (2018)). Since the first large-scale molecular phylogenetic reconstruction of the Polygonaceae in 2003 (Lamb-Frye and Kron 2003), the infrafamilial relationships have gradually been resolved in subsequent studies (e.g. Kim and Donoghue (2008a, b); Kim et al. (2008); Sanchez and Kron (2008, 2009, 2011); Galasso et al. (2009); Sanchez et al. (2009, 2011); Burke et al. (2010); Tavakkoli et al. (2010, 2015); Yurtseva et al. (2010, 2016); Schuster et al. (2011a, b, 2015); Kempton (2012)) and its classification at subfamilial and tribal levels has been significantly improved (Sanchez and Kron 2008; Galasso et al. 2009; Sanchez et al. 2009, 2011; Schuster et al. 2011b, 2015). The majority of genera have been included in previous molecular phylogenetics and their monophyly and circumscription were validated, but a few genera were recircumscribed, such as Atraphaxis L., Koenigia L., Polygonum L., Ruprechtia C.A.Mey. etc. As a result, some new genera were erected, i.e. Duma T.M.Schuster (Schuster et al. 2011b), Salta Adr.Sanchez and Magoniella Adr.Sanchez (Sanchez and Kron 2011), Bactria O.V.Yurtseva & E.V.Mavrodiev (Yurtseva et al. 2016), Persepolium O.V.Yurtseva & E.V.Mavrodiev (Yurtseva et al. 2017) and several old genera have been reduced, for example, Aconogonon (Meisn.) Rchb., Rubrivena M.Král and Emex Neck. ex Campd. (Schuster et al. 2015), Parapteropyrum A.J.Li (Sanchez et al. 2011), Polygonella Michx. (Schuster et al. 2011a) etc. However, due to a dearth of materials or insufficient molecular data to date, the systematic positions of two resistant genera, *Harpagocarpus* Hutch. & Dandy and Eskemukerjea Malick & Sengupta, have not yet been thoroughly evaluated in molecular analyses (Schuster et al. 2015).

The genus *Harpagocarpus* was established on the basis of its distinct fruit morphology (Hutchinson and Dandy 1926) and contains the sole species, *H. snowdenii* Hutch. & Dandy, which was originally recorded only in Uganda, but now has been reported from Kenya, Tanzania, Rwanda and Cameroon (Ayodele 2003). Jacques-Félix (1946)

described *Fagopyrum ciliatum* Jacq.-Fél. from Cameroon, but according to Graham (1958), it is merely a synonym of *H. snowdenii*. Due to its unique appurtenances growing along the achene ribs, which are long purple setae with the radially arranged retrorse barbs at the tip of each seta (Fig. 1), *H. snowdenii* is a distinctive species in Polygonaceae (Hutchinson and Dandy 1926).

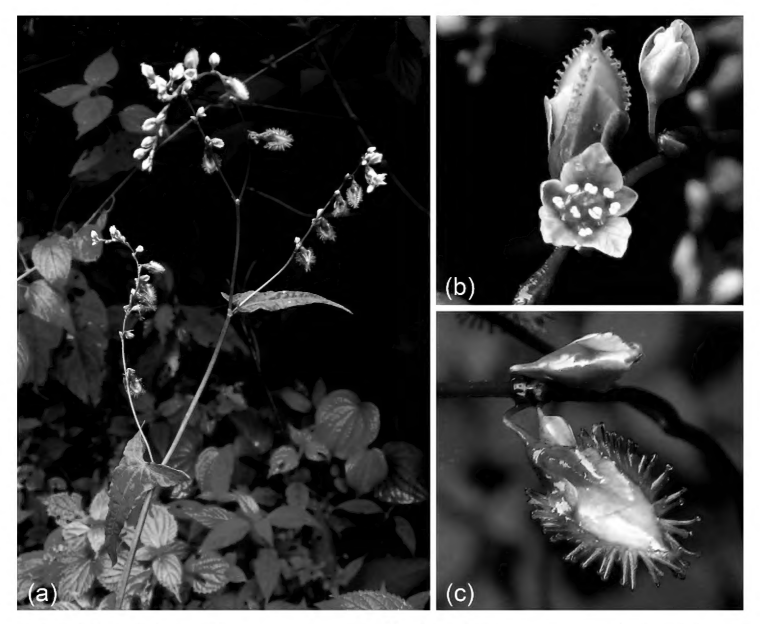
In the protologue, *Harpagocarpus* was morphologically compared to *Polygonum* L. and *Fagopyrum* Mill., but it was thought to resemble the latter considerably more on its broad cotyledons, large and obviously exerted fruits and the shape and venation of the leaves (Hutchinson and Dandy 1926). On the basis of anatomical similarities, Haraldson (1978) hypothesised that *Harpagocarpus* may be closely related to *Fallopia* Adans. However, Ronse Decraene and Akeroyd (1988) argued against this hypothesis and pointed out that *Harpagocarpus* and *Fagopyrum* share considerable similarity in the morphology of floral characteristics. Hong (1988) further reduced *Harpagocarpus* to a synonym of *Fagopyrum* and proposed the new combination *F. snowdenii* (Hutch. & Dandy) S.P.Hong for *H. snowdenii* after concluding from additional palynological research. Though this treatment has been followed in some literature (e.g. Brandbyge (1993); Friis and Vollesen (1998); Sanchez et al. (2011); de Klerk et al. (2015)), it was, nonetheless, recommended that molecular data be used to confirm the phylogenetic position of *Harpagocarpus* (Schuster et al. 2015).

In the present study, we obtained a few precious pieces of leaf materials of *H. snowdenii* from the specimen *Marshall A.R. WK 374* (detailed information available from: http://legacy.tropicos.org/image/100427626), which provided us an invaluable opportunity to investigate the phylogenetic position of *Harpagocarpus*, based on additional molecular data. We adopted two steps of phylogenetic analyses to infer the generic and specific affinities of *H. snowdenii*. Firstly, we used three chloroplast DNA (cpDNA) markers (*matK*, *rbcL* and *trnL-F*) to present the backbone phylogeny of Polygonaceae and affirmed the position of *Harpagocarpus* in *Fagopyrum*. Subsequently, based on five cpDNA regions (*accD*, *matK*, *psbA-trnH*, *rbcL* and *trnL-F*), we further reconstructed the phylogeny of *Fagopyrum* and clarified the accurate specific relationships of *F. snowdenii* within *Fagopyrum*.

#### Materials and methods

# Taxon sampling, choice of markers and datasets

We employed *matK*, *rbcL* and *trnL-F* sequences, which have been extensively used in previous studies (e.g. Lamb-Frye and Kron (2003); Sanchez and Kron (2008); Sanchez et al. (2009, 2011); Burke et al. (2010); Schuster et al. (2015)), to generate a concatenated cpDNA dataset (D1) for reconstructing the backbone phylogeny of Polygonaceae. The ingroup taxa were selected from the entire family to cover all recognised tribal clades (Sanchez et al. 2011; Kempton 2012; Schuster et al. 2015) with at least one representative of each genus. A total of 37 genera and 77 species were sampled. *Plumbago auriculata* Lam. from Plumbaginaceae, which is the sister family of Polygonaceae (Yao et al. 2019;



**Figure 1.** Morphology of *Harpagocarpus snowdenii* Hutch. & Dandy **a** ovate-triangular leaf blades and raceme-like inflorescences **b** an opened flower and a young fruit with minute papillae **c** a mature fruit with long purple setae. (Photographer: Vincent Droissart).

Li et al. 2021), was selected as the outgroup taxon. The source publications or voucher information for all sequences were listed in Suppl. material 1: table S1.

As the analyses of the D1 dataset demonstrated that *Harpagocarpus* is nested within *Fagopyrum*, we designed another dataset (D2) using five cpDNA regions (*accD*, *matK*, *psbA-trnH*, *rbcL* and *trnL-F*), with an expanded sampling of *Fagopyrum* aiming for a more accurate placement of *H. snowdenii* (= *F. snowdenii*). The ingroups of D2 dataset included 33 taxa of *Fagopyrum* covering most of the recognised species in the genus and the outgroup taxon was set as *Pteroxygonum giraldii* Damm. et Diels according to the results presented in Schuster et al. (2015). Voucher information and GenBank accession numbers for taxa used in the D2 dataset are provided in Suppl. material 1: table S2.

# DNA extraction, amplification and sequencing

Total genomic DNA was extracted from fresh or silica gel dried leaves following the manufacturer's specifications of the DNEasy Plant Mini Kit (Qiagen, Valencia, CA, USA). After extraction, the DNA was resuspended in double-distilled water and kept

at -40 °C for polymerase chain reaction (PCR). The PCR reactions and amplification protocol followed Schuster et al. (2011a). The amplified products were purified using a PCR Product Purification Kit (Shanghai SBS, Biotech Ltd., China). Sequencing reactions were conducted with the forward and reverse PCR primers using the DYEnamic ET Terminator Cycle Sequencing Kit (Amersham Biosciences, Little Chalfont, Buckinghamshire, U.K.) with an ABI PRISM 3730 automatic DNA sequencer (Shanghai Sangon Biological Engineering Technology & Services Co., Ltd., Shanghai, China). Both strands of the DNA were sequenced with overlapping regions to ensure that each base was unambiguous. Electropherograms were assembled and consensus sequences were generated with Geneious Prime 2022.0.2 platform.

## Phylogenetic analysis

Sequencher version 5.4.6 (Gene Codes Corporation 2021) was used to evaluate chromatograms for base confirmation and editing contiguous sequences. All DNA sequences were initially aligned using Clustal X version 2.1 (Larkin et al. 2007) and adjusted manually in BioEdit Sequence Alignment Editor version 7.2.1 (Hall 1999).

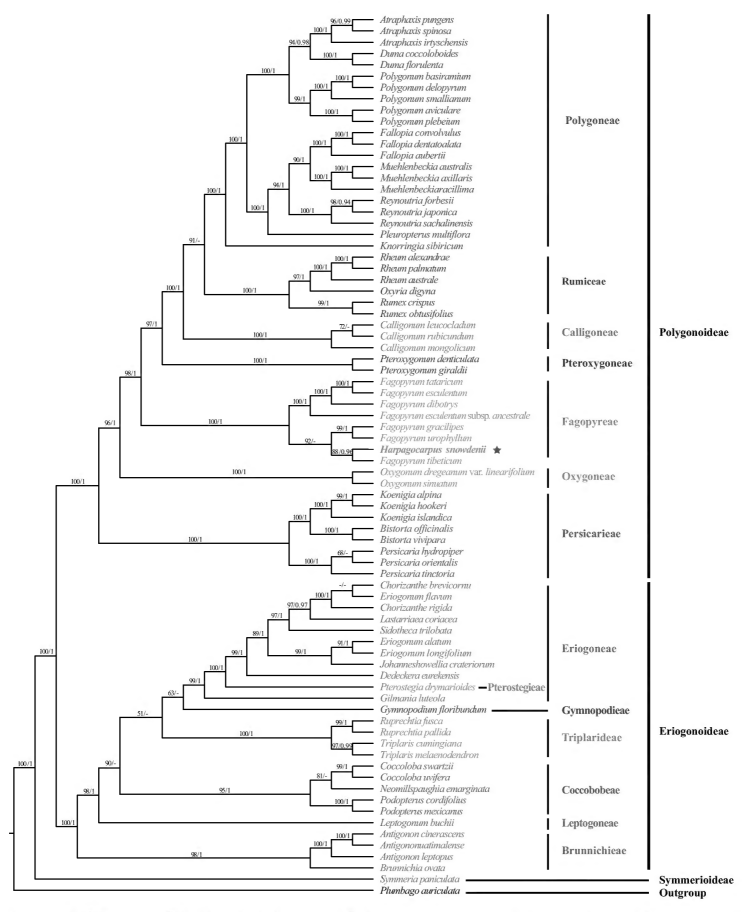
Phylogenetic analyses were conducted, based on the combined cpDNA dataset D1 and D2. The cpDNA regions were supposedly safe to be combined in phylogenetic analyses (Olmstead and Sweere 1994) because the plastid genome is mostly uniparentally inherited (Soltis and Soltis 1998). The datasets were analysed separately using the methods of Maximum Likelihood (ML) and Bayesian Inference (BI).

ML and BI analyses were carried out using RAxML-HPC2 version 8.2.9 (Stamatakis 2014) and MrBayes version 3.2.2 (Ronquist et al. 2012) as implemented on the CIPRES Science Gateway (Miller et al. 2010), respectively. The ML analysis was performed under the GTRGAMMA model with the bootstrap iterations (-# | -N) set to 1000. The BI analysis was executed with most of the default parameters, but manually setting the following: the best substitution types (Nst) and rate distribution models (rates) that were determined by the jModelTest version 2.1.7 (Darriba et al. 2012), sampling one tree every 3000 generations for 100 million generations, stop early if the convergence diagnostic falls below the stop value 0.001 and show tree probabilities on the 50% majority-rule consensus tree with simple output format.

#### Results

# Phylogenetic analyses of Polygonaceae

The concatenated cpDNA dataset D1 has 78 aligned sequences and comprises 4167 characters (1585 bp for *matK*, 1432 bp for *rbcL* and 1150 bp *trnL-F*, respectively), of which 1756 are variable (42.14%) and 1181 are parsimony-informative (28.34%). The ML and BI analyses, based on dataset D1, generated nearly identical topologies (Suppl. material 1: figs S1, S2); therefore, only the ML tree is presented, with ML bootstrap (BS) and posterior probabilities (PP) values marked on each branch, respectively (Fig. 2).



**Figure 2.** Maximum Likelihood phylogram of Polygonaceae as inferred from analysis of the combined cpDNA dataset of matK, rbcL and trnL-F. Support values  $\geq 50\%$  BS or 0.90 PP are displayed above the branches, respectively. The tribal classification of Eriogonoideae followed Sanchez and Kron (2008) and Kempton (2012) and that of Polygoniodeae followed Sanchez et al. (2011) and Schuster et al. (2015). The green star indicates the position of Harpagocarpus snowdenii.

The ingroup (Polygonaceae) is well supported as monophyletic (Fig. 2; BS = 100%, PP = 1.00; all support values follow this order hereafter). Within Polygonaceae, the first branch, represented by *Symmeria paniculata* Benth., is Symmerioideae which is

sister to a large clade comprising Eriogonoideae and Polygonoideae. Within Eriogonoideae, six tribes are recovered with Brunnichieae emerging as the first divergent clade and then subsequently followed by Leptogoneae, Coccobobeae, Triplarideae, Gymnopodieae and Eriogoneae+Pterostegieae. *Pterostegia drymarioides* Fisch. & C.A.Mey. of Pterostegieae is shown to be nested within Eriogoneae in our analyses. In Polygonoideae, all seven tribes are fully supported as monophyletic (Fig. 2) with Persicarieae, Oxygoneae, Fagopyreae, Pteroxygoneae, Calligoneae and Rumiceae successively sister to the rest. With the inclusion of *Harpagocarpus*, *Fagopyrum* obtained high support values (Fig. 2; 100, 1.00).

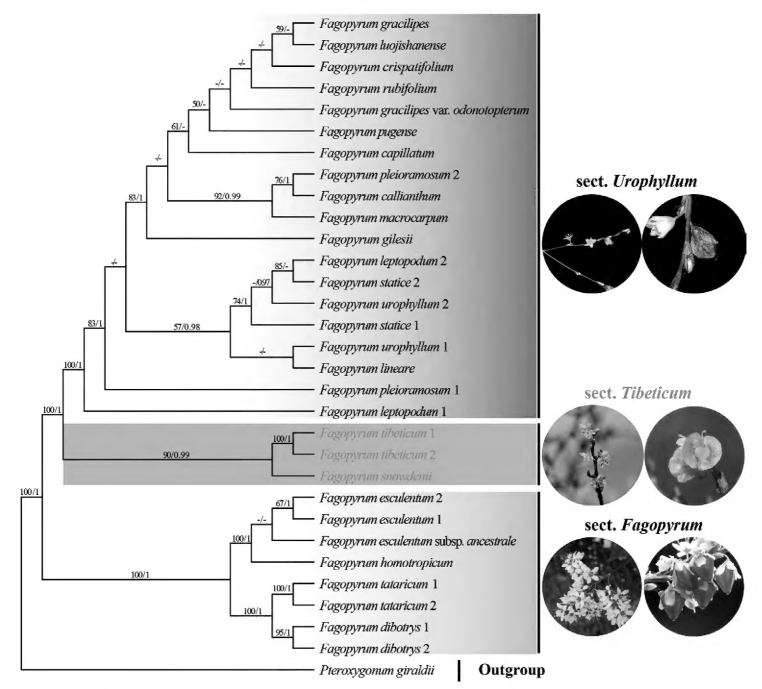
## Phylogenetic analyses of Fagopyrum

The combined dataset D2 has 31 aligned sequences and comprises 6378 characters (1425 bp for accD, 2278 bp for matK, 513 bp for psbA-trnH, 1278 bp for rbcL and 883 bp for trnL-F), of which 735 are variable (11.52%) and 428 are parsimony-informative (6.71%). ML and BI trees generated from the D2 dataset yielded similar topologies (Suppl. material 1: figs S3, S4); thus, only the ML tree is shown (Fig. 3). In both of the analyses, the monophyly of Fagopyrum was strongly supported and three monophyletic subclades were recovered: the first subclade comprises F esculentum Moench, F esculentum subsp. ancestrale Ohnishi, F homotropicum Ohnishi, F tataricum (L.) Gaertn. and F dibotrys (D.Don) H.Hara (100, 1.00), the second one is formed by F snowdenii ( $\equiv Harpagocarpus snowdenii$ ) and F tibeticum (A.J.Li) Adr.Sanchez & Jan.M.Burke (90, 0.99) and the third includes the remaining taxa of the genus.

#### Discussion

# Phylogenetic placement of Harpagocarpus in Polygonaceae

After 20 years of molecular reconstruction of Polygonaceae (e.g. Kim and Donoghue (2008a); Sanchez and Kron (2008, 2009, 2011); Galasso et al. (2009); Sanchez et al. (2009, 2011); Burke et al. (2010); Tavakkoli et al. (2010, 2015); Yurtseva et al. (2010, 2016, 2017); Schuster et al. (2011a, b, 2015); Tian et al. (2011); Kempton (2012); Fan et al. (2021)), only a few recalcitrant genera, such as *Harpagocarpus* and *Eskemukerjea*, have not been included in molecular analyses and their phylogenetic positions are still unresolved. *Harpagocarpus* is distinct from all other genera of Polygonaceae in having a remarkable fruit that has long setae growing along the ribs with the radially arranged retrorse barbs at the tip of each seta (Hutchinson and Dandy 1926) (Fig. 1). The current study is the first to include the genus *Harpagocarpus* in molecular phylogenetics and it demonstrates in all analyses, based on the cpDNA datasets D1 and D2, that *H. snowdenii*, the sole species of the genus, is deeply nested within *Fagopyrum* (Figs 2, 3). The additional molecular evidence undoubtedly confirms the previous hypothesis that *Harpagocarpus* and *Fagopyrum* are congeneric in respect of



**Figure 3.** Maximum Likelihood phylogram of *Fagopyrum* as inferred from analysis of the combined cpDNA dataset of accD, matK, psbA-trnH, rbcL and trnL-F. Support values  $\geq 50\%$  BS or 0.90 PP are displayed above the branches, respectively. The two black boxes covered the cymosum group and the urophyllum group as defined in Yasui and Ohnishi (1998a), respectively. The green box indicates the third group, namely the tibeticum group, as revealed in the present study. The representative photos in the circles showing the inflorescences and the fruits of urophyllum group, tibeticum group and cymosum group are E urophyllum, E tibeticum and E dibotrys, respectively. All photos were taken by Bo Li.

morphological, anatomical and palynological investigations (Hong 1988; Ronse Decraene and Akeroyd 1988; Hong et al. 1998).

Morphologically, *H. snowdenii* has sagittate to ovate-triangular leaf blades, seven palmate veins, large and clearly exerted fruits from the persistent tepals and broad cotyledons, which are very similar to those traits presented in *Fagopyrum* species (Hutchinson and Dandy 1926). Anatomical studies showed that *H. snowdenii* and *Fagopyrum* species share a series of similar floral characteristics, such as nectaries which are present as receptacular mamillae behind the stamens, inner stamens which are always linked with two lateral nectaries and cells of the inner tepal epidermis which are rectangular

to elongated (Hong 1988; Ronse Decraene and Akeroyd 1988). Palynologically, Ronse Decraene and Akeroyd (1988) emphasised that they observed an identical pollen structure between *Harpagocarpus* and *Fagopyrum* and Hong (1988) further noted that it is hardly possible to find any pollen morphological differences between *Harpagocarpus* and the species of *Fagopyrum*. The pollen of *Fagopyrum* is ovate, tricolpate with narrow furrows and a reticulate surface pattern. The pollen grains of *Harpagocarpus* are slightly smaller than those of *Fagopyrum* species, but they undoubtedly belong to the same pollen type (Hong 1988; Ronse Decraene and Akeroyd 1988). Considering this evidence, Ronse Decraene and Akeroyd (1988) suggested that *H. snowdenii* should probably be included within *Fagopyrum*, perhaps as a separate section and Hong (1988) formally combined *H. snowdenii* as *F. snowdenii*.

It is noteworthy to point out that our molecular analyses not only supported the amalgamation of *Harpagocarpus* with *Fagopyrum*, but also clarified the accurate specific relationships of F. snowdenii within Fagopyrum, which was stably supported to be a sister of F. tibeticum using cpDNA sequences (Figs 2, 3). F. tibeticum was originally described in the monotypic genus *Parapteropyrum* A.J.Li as *P. tibeticum* A.J.Li, which is a shrub, endemic to the central Qinghai-Tibetan Plateau of China and is characterised by having fascicled leaves, terminal raceme-like inflorescences, five unequally lobed tepals with the outer two smaller, perianth persistent and accrescent in fruit, three free styles with capitate stigmas and trigonous achenes with broad wings along ribs (Li 1981). P. tibeticum was considered to be most similar to Pteropyrum Jaub. & Spach in gross morphology (Li 1981), but surprisingly tested to be a member of Fagopyrum in molecular phylogenetic studies (Sanchez et al. 2009; Tavakkoli et al. 2010; Tian et al. 2011) and, thus, formally combined in *Fagopyrum* as *F. tibeticum* (Sanchez et al. 2011). The origin of the woody *F. tibeticum* was thought to be a consequence of the large-scale uplift of the Qinghai-Tibetan Plateau which not only promoted continental species radiation, but also the secondary feature of woodiness in a few herbaceous lineages in response to strong selection pressures (Tian et al. 2011).

The inclusion of *F. tibeticum* in *Fagopyrum* has updated our knowledge of morphology in the genus, but now, the sister relationships between *F. snowdenii* and *F. tibeticum*, revealed in our molecular analyses, would not only further expand the morphological variation of *Fagopyrum*, but also shed light on the thinking of the biogeographical origin of the genus, because *F. snowdenii* is the only species of *Fagopyrum* distributed in Africa, while all other congeneric taxa occur mainly in East Asia. Jacques-Félix (1946) suggested that *Fagopyrum* perhaps entered Africa via a Middle Asian pathway during the Quaternary-periglacial period, just like other genera with both Afromontane and Central Asian representatives, such as *Cicer* L. and *Colutea* L. (Chapman and White 1970). However, de Klerk et al. (2015) stated that long-distance transport of pollen grains of *F. snowdenii* from Asia to Africa seems unlikely, but alternatively, they found out there are indications from pollen and macrofossils that a wild *Fagopyrum* ancestor may have been widespread in western Eurasia during the Late Tertiary and the Pleistocene Ice-Ages and became extinct afterwards. *F. snowdenii* may represent the only surviving African lineage that split from the wild widespread *Fagopyrum* ancestor.

## Infrageneric relationships within Fagopyrum

Fagopyrum is a small genus comprised of ca. 25 species according to the most updated classification (Ohsako and Li 2020). The genus is economically important and well known for containing two domesticated common buckwheat, i.e. F. esculentum and F. tataricum which have been widely cultivated in Australia, Asia, Europe and North America for producing gluten-free grains (Li and Hong 2003). Geographically, most of the wild species of Fagopyrum are mainly distributed in mountainous regions of southwest China, a few are endemic to the south-eastern edge of the Qinghai-Tibetan Plateau (Ohnishi and Matsuoka 1996; Ohnishi 1998; Li and Hong 2003) and only the F. snowdenii confirmed in the present study is occurring in Africa (Hutchinson and Dandy 1926; Ayodele 2003). Eastern Tibet to western Sichuan of China was indicated to be the birthplace of the two cultivated common buckwheat in the AFLP (amplified fragment length polymorphism) analysis (Konishi et al. 2005). Taxonomically, Fagopyrum was separated from the large and heterogenous Linnaeus's genus Polygonum L. (Miller 1754) and has long been treated as a section of *Polygonum* (e.g. Meisner (1856); Samuelsson (1929); Steward (1930)) or considered to be an independent genus, but closely related to *Polygonum* (e.g. Dammer (1894); Gross (1913); Hedberg (1946); Haraldson (1978); Ronse Decraene and Akeroyd (1988)). In the context of the molecular phylogeny of Polygonaceae, Fagopyrum was not only supported as a monophyletic genus, but also indicated to represent an isolated tribal clade in the subfamily Polygonoideae (Sanchez et al. 2011; Schuster et al. 2015). Morphologically and anatomically, Fagopyrum could be distinguished from other genera of Polygonoideae by having large conduplicate cotyledons and/or embryos in the central region in achene (Dammer 1894; Gross 1913; Nakai 1926; Chapman and White 1970; Sanchez et al. 2011).

Within Fagopyrum, two groups have been recognised in classical taxonomy, based on the morphology of inflorescence and the achene size: one group was mainly represented by *F. cymosum* (Trevir.) Meisn. (= *F. dibotrys*), *F. esculentum* and *F. tataricum* and characterised by having corymbose inflorescences with many branching and dense flowers and the achene greatly exceeding the perianth, while the other group is composed of other species (including F. urophyllum (Bureau & Franch.) H.Gross) having racemelike inflorescences with sparse flowers and the achene completely enclosed in perianth (Gross 1913; Roberty and Vautier 1964; Ohnishi and Matsuoka 1996) (Fig. 3). These two groups are mostly concordant with the cymosum group and the urophyllum group defined by Yasui and Ohnishi (1998a) in molecular phylogenetic analyses using DNA sequences of the nuclear internal transcribed spacer (nrITS) and cpDNA region rbcLaccD. Other molecular studies, no matter using isozyme variability and RFLP (Ohnishi and Matsuoka 1996), cpDNA sequences (Yasui et al. 1998; Ohsako et al. 2001; Jin et al. 2018), nuclear genes or regions (Yasui and Ohnishi 1998b; Nishimoto et al. 2003) and complete plastomes (Fan et al. 2021; Li et al. 2022), all clearly indicated that the cymosum group and the urophyllum group are both monophyletic clades.

In our present analyses, the above-mentioned two clades were recovered too, but the third clade, formed by *F. snowdenii* and *F. tibeticum*, was discovered, which is

sister to the 'Urophyllum' clade (Fig. 3). We failed to generate any nuclear sequences from the specimen sample of *F. snowdenii*; thus, we could not test the sister relationships between *F. snowdenii* and *F. tibeticum*, as well as the sister relationships between F. snowdenii +F. tibeticum clade and the 'Urophyllum' clade in nuclear analysis. However, when only F. tibeticum was included in the ITS analysis, the topology of the phylogenetic tree is similar to that yielded from the combined cpDNA dataset, in which F. tibeticum is sister to the 'Urophyllum' clade clade (Tian et al. 2011). Considering the sister relationships between F. snowdenii and F. tibeticum could be additionally supported by morphological and palynological evidence, such as racemelike inflorescences, unequal tepals with the outer two smaller, perianth accrescent in fruit, large achenes greatly exceeding the perianth, special appurtenances (either wings or setae) growing along the fruit ribs and smaller pollen grains than the other Fagopyrum species (Hutchinson and Dandy 1929; Ronse Decraene and Akeroyd 1988; Hong 1995), we believe that F. snowdenii and F. tibeticum represent a separate clade in *Fagopyrum*. Future analyses, based on more comprehensive sampling and using nuclear sequences data, may further confirm or update the infrageneric relationships of Fagopyrum as inferred in this study. As far as the current results are concerned, a sectional classification for Fagopyrum is here proposed, based on the differentiation of gross morphology in the three clades, which is the first infrageneric classification of the genus.

#### Taxonomic treatment

#### Fagopyrum Mill., Gard. Dict. Abr.

- Fagopyrum Mill., Gard. Dict. Abr., ed. 4, 495. 1754 [≡ Polygonum sect. Fagopyrum (Mill.) Meisn., Monogr. Polyg. 43, 61. 1826.] Type: Fagopyrum esculentum Moench (≡ Polygonum fagopyrum L.).
- = Harpagocarpus Hutch. & Dandy, Bull. Misc. Inform. Kew. 364. 1926 Type: Harpagocarpus snowdenii Hutch. & Dandy [≡ Fagopyrum snowdenii (Hutch. & Dandy) S.P.Hong].
- = Parapteropyrum A.J.Li, Acta Phytotax. Sin. 19: 330. 1981 Type: Parapteropyrum tibeticum A.J.Li [≡ Fagopyrum tibeticum (A.J.Li) Adr.Sanchez & Jan.Burke].

## Fagopyrum sect. Fagopyrum

**Type.** *Fagopyrum esculentum* Moench. ( $\equiv$  *Polygonum fagopyrum* L.).

**Diagnosis.** This section is characterised by having large corymbose inflorescences with many branches and dense flowers and large achenes greatly exceeding the persistent perianth.

**Species.** F. dibotrys, F. esculentum subsp. esculentum, F. esculentum subsp. ancestrale, F. homotropicum and F. tataricum.

**Distribution.** Bhutan, India, Myanmar, Nepal, Pakistan, Thailand and Vietnam of southern and south-eastern Asia and southern and south-western China.

## Fagopyrum sect. Tibeticum Bo Li & M.L.Zhou, sect. nov.

urn:lsid:ipni.org:names:77315008-1

**Type.** Fagopyrum tibeticum (A.J.Li) Adr. Sanchez & Jan. Burke (≡ Parapteropyrum tibeticum A.J.Li).

**Diagnosis.** The new section is characterised by having raceme-like inflorescences with sparse flowers, large achenes with appurtenances (wings or setae) along the ribs and greatly exceeding the perianth and persistent perianth accrescent in fruit.

**Species.** *F. snowdenii* and *F. tibeticum*.

**Distribution.** Cameroon, Kenya, Rwanda, Tanzania and Uganda of Africa (*F. snowdenii*) and Tibet of south-western China (*F. tibeticum*).

## Fagopyrum sect. Urophyllum Bo Li & M.L.Zhou, sect. nov.

urn:lsid:ipni.org:names:77315009-1

**Type.** Fagopyrum urophyllum (Bureau & Franch.) H.Gross (≡ Polygonum urophyllum Bureau & Franch.).

**Diagnosis.** This new section is characterised by having raceme-like, spicate, capitate or paniculate inflorescences with mostly sparse or rarely dense flowers and achenes completely enclosed in the persistent perianth.

**Species.** F. callianthum Ohnishi, F. capillatum Ohnishi, F. caudatum (Sam.) A.J.Li, F. crispatifolium J.L.Liu, F. densovillosum J.L.Liu, F. gilesii (Hemsl.) Hedberg, F. gracilipedoides Ohsako & Ohnishi, F. gracilipes (Hemsl.) Dammer, F. jinshaense Ohsako & Ohnishi, F. leptopodum (Diels) Hedberg var. leptopodum, F. leptopodum var. grossii (Lévl.) Lauener & D.K.Ferguson, F. lineare (Sam.) Haraldson, F. longistylum M.L.Zhou & Y.Tang, F. longzhoushanense J.R.Shao, F. luojishanense J.R.Shao, F. macrocarpum Ohsako & Ohnishi, F. pleioramosum Ohnishi, F. pugense Y.Tang, F. qiangcai D.Q.Bai, F. rubifolium Ohsako & Ohnishi, F. statice H.Gross, F. urophyllum (Bureau & Franch.) H.Gross, F. wenchuanense J.R.Shao.

Distribution. Guizhou, Sichuan and Yunnan Provinces of southwest China.

# Identification keys to three sections of Fagopyrum

## **Acknowledgements**

The authors are grateful to administrators of the Missouri Botanical Garden Herbarium (MO) for providing the specimen sample of *Harpagocarpus snowdenii*, to Dr. Tanja M. Schuster in the Naturhistorisches Museum Wien for sharing molecular data of Polygonaceae and to Vincent Droissart in the Institut de Recherche pour le Développement (IRD, France) for providing field photos of *H. snowdenii*. This study was jointly supported by the National Key R&D Program of China (2021YFD1200100/2021YFD1200105) and the National Natural Science Foundation of China (32160047, 31900181, 32161143005).

## References

- Ayodele AE (2003) The distribution and ecology of the family Polygonaceae in West Africa. Nigerian Journal of Ecology 5: 46–49.
- Brandbyge J (1993) Polygonaceae. In: Kubitzki K (Ed.) The Families and Genera of Vascular Plants (Vol. 6). Springer, Berlin, 531–544. https://doi.org/10.1002/fedr.19921030317
- Brockington SF, Alexandre R, Ramdial J, Moore MJ, Crawley S, Dhingra A, Hilu K, Soltis DE, Soltis PS (2009) Phylogeny of the Caryophyllales sensu lato: Revisiting hypotheses on pollination biology and perianth differentiation in the core Caryophyllales. International Journal of Plant Sciences 170(5): 627–643. https://doi.org/10.1086/597785
- Burke JM, Sanchez A, Kron KA, Luckow M (2010) Placing the woody tropical genera of Polygonaceae: A hypothesis of character evolution and phylogeny. American Journal of Botany 97(8): 1377–1390. https://doi.org/10.3732/ajb.1000022
- Chapman JD, White F (1970) The Evergreen Forests of Malawi. Oxford University Press, Oxford, 190 pp. https://doi.org/10.1017/S0030605300011777
- Cuénoud P, Savolainen V, Chatrou LW, Powell M, Grayer RJ, Chase MW (2002) Molecular phylogenetics of Caryophyllales based on nuclear 18S rDNA and plastid *rbcL*, *atpB*, and *matK* DNA sequences. American Journal of Botany 89(1): 132–144. https://doi.org/10.3732/ajb.89.1.132
- Dammer U (1894) Polygonaceae. In: Engler A, Prantl K (Eds) Die Natürlichen Pflanzenfamilien (Vol. 3). Verlag von Wilhelm Engelmann, Leipzig, 36 pp.
- Darriba D, Taboada GL, Doallo R, Posada D (2012) jModelTest 2: More models, new heuristics and parallel computing. Nature Methods 9(8): e772. https://doi.org/10.1038/nmeth.2109
- de Klerk P, Couwenberg J, Joosten H (2015) Pollen and macrofossils attributable to *Fagopyrum* in western Eurasia prior to the Late Medieval: An intercontinental mystery. Palaeogeography, Palaeoclimatology, Palaeoecology 440: 1–21. https://doi.org/10.1016/j.palaeo.2015.08.024
- Fan Y, Jin Y, Ding M, Tang Y, Cheng J, Zhang K, Zhou M (2021) The complete chloroplast genome sequence of eight *Fagopyrum* species: Insights into genome evolution and phylogenetic relationships. Frontiers in Plant Science 12: e799904. https://doi.org/10.3389/fpls.2021.799904

- Friis I, Vollesen K (1998) Flora of the Sudan-Uganda Border Area East of the Nile I: Catalogue of Vascular Plants. Munksgaard, Copenhagen, Denmark, 389 pp.
- Galasso G, Banfi E, Mattia FD, Grassi F, Sgorbati S, Labra M (2009) Molecular phylogeny of *Polygonum* L. s.l. (Polygonoideae, Polygonaeae), focusing on European taxa: Preliminary results and systematic considerations based on *rbcL* plastidial sequence data. Atti della Societa Italiana di Scienze Naturali e del Museo Civico di Storia Naturale di Milano 150: 113–148.
- Gene Codes Corporation (2021) Gene Codes Corporation. http://www.genecodes.com [accessed on 22 October 2021]
- Graham RA (1958) Polygonaceae. In: Turill WB, Milne-Redhead E (Eds) Flora of Tropical East Africa (Vol. 1). Crown Agents, London, 40 pp.
- Gross MH (1913) Remarques sur les Polygonés de L'Asie Orientale. Bulletin de Geographie Botanique 23: 7–32.
- Hall TA (1999) BioEdit: A user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. Nucleic Acids Symposium Series 41: 95–98. https://doi.org/10.14601/Phytopathol\_Mediterr-14998u1.29
- Haraldson K (1978) Anatomy and taxonomy in Polygonaceae subfam. Polygonoideae Meissn. [sic.] emend. Jaretzky. Symbolae Botanicae Upsalienses 22: 1–95.
- Hedberg O (1946) Pollen morphology in the genus *Polygonum* L. s. lat. and its taxonomical significance. Svensk Botanisk Tidskrift 40: 371–404.
- Hong SP (1988) A pollen morphological re-evaluation of *Harpagocarpus* and *Eskemukerjea* (Polygonaceae). Grana 27(4): 291–295. https://doi.org/10.1080/00173138809429951
- Hong SP (1995) Pollen morphology of *Parapteropyrum* and some putatively related genera (Polygonaceae-Atraphaxideae). Grana 34(3): 153–159. https://doi.org/10.1080/00173139509429039
- Hong SP, Ronse Decraene LP, Smets E (1998) Systematic significance of tepal surface morphology in tribes Persicarieae and Polygoneae (Polygonaceae). Botanical Journal of the Linnean Society 127(2): 91–116. https://doi.org/10.1111/j.1095-8339.1998.tb02091.x
- Hutchinson J, Dandy JE (1926) A new genus from Uganda. Bulletin Miscellaneous Information 8(8): 363–365. https://doi.org/10.2307/4118200
- Jacques-Félix H (1946) Description d'un *Fagopyrum* Africain. Bulletin du Muséum National d'Histoire Naturelle 2: 409–411.
- Jin JW, Li DR, Chen SF, Li B (2018) A common *Bistorta* was misidentified as a novel species in *Fagopyrum* (Polygonaceae): The confirmation of the taxonomic identify of *F. hailuogouense* by morphological and molecular evidences. Phytotaxa 348(3): 221–228. https://doi.org/10.11646/phytotaxa.348.3.5
- Judd WS, Campbell CS, Kellogg EA, Stevens PF, Donoghue MJ (2007) Plant Systematics: A Phylogenetic Approach. Sinauer Associates, Sunderland, 678 pp.
- Kempton EA (2012) Systematics of Eriogonoideae s.s. (Polygonaceae). Systematic Botany 37(3): 723–737. https://doi.org/10.1600/036364412X648698
- Kim ST, Donoghue MJ (2008a) Incongruence between cpDNA and nrITS trees indicates extensive hybridization within *Eupersicaria* (Polygonaceae). American Journal of Botany 95(9): 1122–1135. https://doi.org/10.3732/ajb.0700008

- Kim ST, Donoghue MJ (2008b) Molecular phylogeny of *Persicaria* (Persicarieae, Polygonaceae). Systematic Botany 33(1): 77–86. https://doi.org/10.1600/036364408783887302
- Kim ST, Sultan SE, Donoghue MJ (2008) Allopolyploid speciation in *Persicaria* (Polygonaceae): Insights from a low-copy nuclear region. Proceedings of the National Academy of Sciences of the United States of America 105(34): 12370–12375. https://doi.org/10.1073/pnas.0805141105
- Konishi T, Yasui Y, Ohnishi O (2005) Original birthplace of cultivated common buckwheat inferred from genetic relationships among cultivated populations and natural populations of wild common buckwheat revealed by AFLP analysis. Genes & Genetic Systems 80(2): 113–119. https://doi.org/10.1266/ggs.80.113
- Lamb-Frye AS, Kron KA (2003) *rbcL* phylogeny and character evolution in Polygonaceae. Systematic Botany 28: 326–332. https://doi.org/10.1266/ggs.78.139
- Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, Thompson JD, Gibson TJ, Higgins DG (2007) Clustal W and Clustal X version 2.0. Bioinformatics 23(21): 2947–2948. https://doi.org/10.1093/bioinformatics/btm404
- Li AJ (1981) *Parapteropyrum* A. J. Li: Unum genus novum *Polygonacearum sinicum*. Acta Phytotaxonomica Sinica 19: 330–331.
- Li AR, Hong SP (2003) *Fagopyrum*. In: Wu ZY, Raven PH (Eds) Flora of China: (Ulmaceae through Basellaceae) (Vol. 5). Science Press/Missouri Botanical Garden Press, Beijing / St. Louis, 320–323.
- Li HT, Luo Y, Gan L, Ma PF, Gao LM, Yang JB, Cai J, Gitzendanner MA, Fritsch PW, Zhang T, Jin JJ, Zeng CX, Wang H, Yu WB, Zhang R, van der Bank M, Olmstead RG, Hollingsworth PM, Chase MW, Soltis DE, Soltis PS, Yi T-S, Li D-Z (2021) Plastid phylogenomic insights into relationships of all flowering plant families. BMC Biology 19(1): e232. https://doi.org/10.1186/s12915-021-01166-2
- Li QJ, Liu Y, Wang AH, Chen QF, Wang JM, Peng L, Yang Y (2022) Plastome comparison and phylogenomics of *Fagopyrum* (Polygonaceae): Insights into sequence differences between *Fagopyrum* and its related taxa. BMC Plant Biology 22(1): e339. https://doi.org/10.1186/s12870-022-03715-5
- Meisner CF (1856) Polygonaceae. In: de Candolle A (Ed.) Prodomus Systematis Naturalis Regni Vegetabilis (Vol. 14). Masson, Paris, 185 pp.
- Miller P (1754) The Gardener's Dictionary (Vol. 3). Cambridge University Press, London, 1582 pp.
- Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for Inference of Large Phylogenetic Trees. Proceedings of the Gateway Computing Environments Workshop (GCE). New Orleans, 45–52. https://doi.org/10.1109/GCE.2010.5676129
- Moore MJ, Soltis PS, Bell CD, Burleigh G, Solti DE (2010) Phylogenetic analysis of 83 plastid genes further resolves the early diversification of eudicots. Proceedings of the National Academy of Sciences of the United States of America 107(10): 4623–4628. https://doi.org/10.1073/pnas.0907801107
- Nakai T (1926) Taderui no shinbunruihou (a new classification of Linnaean *Polygonum*). Rigakkai 24: 289–301.

- Nishimoto Y, Ohnishi O, Hasegawa M (2003) Topological incongruence between nuclear and chloroplast DNA trees suggesting hybridization in the *urophyllum* group of the genus *Fagopyrum* (Polygonaceae). Genes & Genetic Systems 78(2): 139–153. https://doi.org/10.1266/ggs.78.139
- Ohnishi O (1998) Search for the wild ancestor of buckwheat I. Description of new *Fagopyrum* (Polygonaceae) species and their distribution in China and Himalayan hills. Fagopyrum 15: 18–28.
- Ohnishi O, Matsuoka Y (1996) Search for the wild ancestor of buckwheat II. Taxonomy of *Fagopyrum* (Polygonaceae) species based on morphology, isozymes and cpDNA variability. Genes & Genetic Systems 71(6): 383–390. https://doi.org/10.1266/ggs.71.383
- Ohsako T, Li C (2020) Classification and systematics of the *Fagopyrum* species. Breeding Science 70(1): 93–100. https://doi.org/10.1270/jsbbs.19028
- Ohsako T, Fukuoka S, Bimb HP, Baniya BK, Yasui Y, Ohnishi O (2001) Phylogenetic analysis of the genus *Fagopyrum* (Polygonaceae), including the Nepali species *F. megacarpum*, based on nucleotide sequence of the *rbcL-accD* region in chloroplast DNA. Fagopyrum 18: 9–14.
- Olmstead RG, Sweere JA (1994) Combining data in phylogenetic systematics: An empirical approach using three molecular data sets in the Solanaceae. Systematic Biology 43(4): 467–481. https://doi.org/10.1093/sysbio/43.4.467
- Roberty G, Vautier S (1964) Les genres de Polygonacées. Boissiera 10: 7–128.
- Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA, Huelsenbeck JP (2012) MrBayes 3.2: Efficient Bayesian phylogenetic inference and model choice across a large model space. Systematic Biology 61(3): 539–542. https://doi.org/10.1093/sysbio/sys029
- Ronse Decraene LP, Akeroyd JR (1988) Generic limits in *Polygonum* and related genera (Polygonaceae) on the basis of floral characters. Botanical Journal of the Linnean Society 98(4): 321–371. https://doi.org/10.1111/j.1095-8339.1988.tb01706.x
- Samuelsson G (1929) Polygonaceae. In: Handel-Mazzetti H (Ed.) Symbolae Sinicae (Vol. 7). Verlag von Julius Springer, Wien, 166–188.
- Sanchez A, Kron KA (2008) Phylogenetics of Polygonaceae with an emphasis on the evolution of Eriogonoideae. Systematic Botany 33(1): 87–96. https://doi.org/10.1600/036364408783887456
- Sanchez A, Kron KA (2009) Phylogenetic relationships of *Afrobrunnichia* Hutch. & Dalziel (Polygonaceae) based on three chloroplast genes and ITS. Taxon 58(3): 781–792. https://doi.org/10.1002/tax.583008
- Sanchez A, Kron KA (2011) Phylogenetic relationships of *Triplaris* and *Ruprechtia*: Re-delimitation of the recognized genera and two new genera for tribe Triplarideae (Polygonaceae). Systematic Botany 36(3): 702–710. https://doi.org/10.1600/036364411X583664
- Sanchez A, Schuster TM, Kron KA (2009) A large-scale phylogeny of Polygonaceae based on molecular data. International Journal of Plant Sciences 170(8): 1044–1055. https://doi.org/10.1086/605121
- Sanchez A, Schuster TM, Burke JM, Kron KA (2011) Taxonomy of Polygonoideae (Polygonaceae): A new tribal classification. Taxon 60(1): 151–160. https://doi.org/10.1002/tax.601013

- Schäferhoff B, Müller KF, Borsch T (2009) Caryophyllales phylogenetics: Disentangling Phytolaccaceae and Molluginaceae and description of Microteaceae as a new isolated family. Willdenowia 39(2): 209–228. https://doi.org/10.3372/wi.39.39201
- Schuster TM, Reveal JL, Kron KA (2011a) Evolutionary relationships within Polygoneae (Polygonaceae: Polygonoideae). Taxon 60: 1653–1666. https://doi.org/10.1002/tax.606010
- Schuster TM, Wilson KL, Kron KA (2011b) Phylogenetic relationships of *Muehlenbeckia*, *Fallopia* and *Reynoutria* (Polygonaceae) investigated with chloroplast and nuclear sequence data. International Journal of Plant Sciences 172(8): 1053–1066. https://doi.org/10.1086/661293
- Schuster TM, Reveal JL, Bayly MJ, Kron KA (2015) An updated molecular phylogeny of Polygonoideae (Polygonaceae): Relationships of *Oxygonum*, *Pteroxygonum*, and *Rumex*, and a new circumscription of *Koenigia*. Taxon 4(6): 1188–1208. https://doi.org/10.12705/646.5
- Soltis PS, Soltis DE (1998) Molecular evolution of 18S ribosomal DNA in angiosperms: Implications for character weighting in phylogenetic analysis. In: Soltis DE, Soltis PS, Doyle JJ (Eds) Molecular Systematics of Plants: DNA Sequencing (Vol. 2). Kluwer, Dordrecht, 188–210. https://doi.org/10.1007/978-1-4615-5419-6\_7
- Stamatakis A (2014) RAxML version 8: A tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics 30(9): 1312–1313. https://doi.org/10.1093/bioinformatics/btu033
- Steward AN (1930) The Polygoneae of Eastern Asia. Contributions from the Gray Herbarium of Harvard University 5(88): 1–129. https://doi.org/10.5962/p.336124
- Tavakkoli S, Kazempour Osaloo S, Maassoumi AA (2010) The phylogeny of *Calligonum* and *Pteropyrum* (Polygonaceae) based on nuclear ribosomal DNA ITS and chloroplast *trnL-F* sequences. Iranian Journal of Biotechnology 8: 7–15.
- Tavakkoli S, Kazempour Osaloo S, Mozaffarian V, Maassoumi AA (2015) Molecular phylogeny of *Atraphaxis* and the woody *Polygonum* species (Polygonaceae): Taxonomic implications based on molecular and morphological evidence. Plant Systematics and Evolution 301(4): 1157–1170. https://doi.org/10.1007/s00606-014-1140-7
- Tian XM, Luo J, Wang AL, Mao KS, Liu JQ (2011) On the origin of the woody buckwheat *Fagopyrum tibeticum* (=*Parapteropyrum tibeticum*) in the Qinghai-Tibetan Plateau. Molecular Phylogenetics and Evolution 61(2): 515–520. https://doi.org/10.1016/j.ympev.2011.07.001
- Walker JF, Yang Y, Feng T, Timoneda A, Mikenas J, Hutchinson V, Edwards C, Wang N, Ahluwalia S, Olivieri J, Walker-Hale N, Majure LC, Puente R, Kadereit G, Lauterbach M, Eggli U, Flores-Olvera H, Ochoterena H, Brockington SF, Moore MJ, Smith SA (2018) From cacti to carnivores: Improved phylotranscriptomic sampling and hierarchical homology inference provide further insight to the evolution of Caryophyllales. American Journal of Botany 105(3): 446–462. https://doi.org/10.1002/ajb2.1069
- Yang Y, Moore MJ, Brockington SF, Soltis DE, Wong GKS, Carpenter EJ, Zhang Y, Chen L, Yan Z, Xie Y, Sage RF, Covshoff S, Hibberd JM, Nelson MN, Smith SA (2015) Dissecting molecular evolution in the highly diverse plant clade Caryophyllales using transcriptome sequencing. Molecular Biology and Evolution 32(8): 2001–2014. https://doi.org/10.1093/molbev/msv081
- Yao G, Jin JJ, Li HT, Yang JB, Mandala VS, Croley M, Mostow R, Douglas NA, Chase MW, Christenhusz MJM, Soltis DE, Soltis PS, Smith SA, Brockington SF, Moore MJ, Yi TS, Li

- DZ (2019) Plastid phylogenomic insights into the evolution of Caryophyllales. Molecular Phylogenetics and Evolution 134: 74–86. https://doi.org/10.1016/j.ympev.2018.12.023
- Yasui Y, Ohnishi O (1998a) Interspecific relationships in *Fagopyrum* (Polygonaceae) revealed by the nucleotide sequences of the *rbcL* and *accD* genes and their intergenic region. American Journal of Botany 85(8): 1134–1142. https://doi.org/10.2307/2446346
- Yasui Y, Ohnishi O (1998b) Phylogenetic relationships among *Fagopyrum* species revealed by the nucleotide sequences of the ITS region of the nuclear rRNA gene. Genes & Genetic Systems 73(4): 201–210. https://doi.org/10.1266/ggs.73.201
- Yasui Y, Ohsako T, Ohnishi O (1998) Evolutionary processes of *Fagopyrum* inferred from the molecular phylogenetic analyses. Proceedings of the 7<sup>th</sup> International Symposium: Current Advances in Buckwheat Research. Winnipeg, Canada, 50–60.
- Yurtseva OV, Troitsky AV, Bobrova VK, Voylokova VN (2010) On taxonomical revision of *Polygonum* s. str. (Polygonaceae): Phylogenetic and morphological data. Botanicheskii Zhurnal 95: 226–247. https://doi.org/10.11646/phytotaxa.314.2.1
- Yurtseva OV, Kuznetsova OI, Mavrodieva ME, Mavrodiev EV (2016) What is *Atraphaxis* L. (Polygonaceae, Polygoneae): Cryptic taxa and resolved taxonomic complexity instead of the formal lumping and the lack of morphological synapomorphies. PeerJ 4: e1977. https://doi.org/10.7717/peerj.1977
- Yurtseva OV, Severova EE, Mavrodiev EV (2017) *Persepolium* (Polygoneae): A new genus in Polygonaceae based on conventional Maximum Parsimony and three-taxon statement analyses of a comprehensive morphological dataset. Phytotaxa 314: 151–194. https://doi.org/10.11646/phytotaxa.314.2.1

# Supplementary material I

## Supplementary information

Authors: Daozhang Min, Wei Shi, Mohammad Mehdi Dehshiri, Yuting Gou, Wei Li, Kaixuan Zhang, Meiliang Zhou, Bo Li

Data type: tables, figures (Pdf file)

Explanation note: Taxa, GenBank accession numbers of DNA sequences with their vouchers or source of publication used in the molecualr dataset and phtlogenetic trees generated from BI and ML analyses.

Copyright notice: This dataset is made available under the Open Database License (http://opendatacommons.org/licenses/odbl/1.0/). The Open Database License (ODbL) is a license agreement intended to allow users to freely share, modify, and use this Dataset while maintaining this same freedom for others, provided that the original source and author(s) are credited.

Link: https://doi.org/10.3897/phytokeys.220.97667.suppl1